## **PROCEEDINGS**

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## ON AN EQUATION FOR THE VISCOSITY OF MIXTURES

By S. K. Charrabertty and P. B. Ganguly

CHEMICAL LABORATORY, SCIENCE COLLEGE, PATNA

Received October 28, 1937

#### SUMMARY

An expression of the form  $\log \eta_{sol} - \frac{1}{3} \log \rho_{sol} = C_1 + C_2 x_2$  can be deduced from Andrade's equation for viscosity on the basis of certain assumptions. The applicability of the above equation has been tested by considering 18 binary systems. On plotting the value of  $\log \eta_{sol} - \frac{1}{3} \log \rho_{sol}$  against  $x_2$  the graphs have been found to be straight lines in many cases. In the case of systems, the components of which have polar properties and are dissimilar in nature, the divergence is marked. This is explained as due to loose compound formation. The viscosity values calculated from MacLeod's equation have been compared with those obtained from the above equation. The latter equation gives somewhat better values.

Frequent attempts have been made to find a relationship between the viscosity of binary mixtures and their compositions. A linear formula based on the mixture law has often been tried but has been found inadequate. It was suggested that the divergence from the linear equation was due to the lack of the correct method of expressing the composition. According to Drucker and Kassel, concentration should be expressed as weight percentages, whilst Kendall considers it more logical to express the composition as molar concentrations. MaeFarlane and Wright have, however, shown that whatever method of plotting the experimental data be adopted, viscosity cannot be expressed as an additive property.

Findlay<sup>5</sup> considers that fluidity, which represents the inverse of viscosity, might be expressed more consistently by the mixture law formula. Bingham and

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Findlay<sup>5</sup> considers that fluidity, which represents the inverse of viscosity, might be expressed more consistently by the mixture law formula. Bingham and

MacMaster3 attempted to express fluidities by an additive formula but found it inadequate.

Attempts have also been made to express viscosity by means of a logarithmic formula. The equation proposed by Arrhenius<sup>2</sup> gave divergent values in concentrated solutions. Kendall modified the Arrhenius formula in which he used molar percentages in place of volume percentages. Later Kendall and Monroe<sup>8</sup> proposed the equation

$$\log \eta_{\text{sol}} = x_1 \log \eta_1 + x_2 \log \eta_2$$

where  $x_1$  and  $x_2$  represented the molar percentages of the two components. Both these equations were subsequently found by them as unsatisfactory.

In a previous paper<sup>6</sup> we have found that the logarithms of the relative viscosities of solutions when plotted against the molar fraction of the solute generally gave straight lines up to 50% mixtures. At higher concentrations the plot was still a straight line but with a different slope. For dilute solutions the formula  $\log \eta/\eta_1 = C_1 x_2 + C_2$ , where  $C_1$  and  $C_2$  were constants, was fairly satisfactory.

Kendall<sup>7</sup> has classified binary mixtures into two groups, the ideal and the non-ideal solutions. A binary system was considered ideal when on mixing the components there was no volume change and no heat evolution. Ideal solutions, however, are hardly attained in practice and even the systems which have been considered by Kendall as ideal show distinct variations in density on mixing. It is evident, therefore, that an allowance has to be made for the change in volume that sets in when the mixture is made. In the following pages an equation involving the necessary density correction has been derived from Andrade's viscosity equation on the basis of certain assumptions.

According to Andrade<sup>1</sup> the viscosity of a pure liquid is given by  $\eta v^{\frac{1}{2}}$   $= A. \ e^{c/vT} \text{ where } v \text{ is the specific volume, } T \text{ the absolute temperature and } A \text{ and } C$ are constants. For a particular temperature the equation can be written as  $\eta/\rho^{\frac{1}{2}}$   $= A. \ e^{\beta\rho} \text{ where } \beta = \frac{C}{T}.$ 

If we assume, as has been done by Spells,  $^{11}$  that Andrade's equation can be applied to the case of solutions also, the viscosity of a solution will be expressed by the equation  $\eta/\rho^{\frac{1}{3}} = A_1$ .  $e^{B\rho}$  where  $A_1$  and B are constants. Expressing in logarithmic form we get

$$\log \eta_{\text{sol}} - \frac{1}{3} \log \rho_{\text{sol}} = C_1 + B\rho_{\text{sol}}$$

where C<sub>1</sub> and B are constants.

The variation of the constant B with concentration has been considered by Spells, who has concluded that the constant is a function of the composition of the mixture. If we assume that Be<sub>sol</sub> is directly proportional to  $x_2$ , where  $x_2$  represents the molar fraction of the solute, the above equation reduces to the form

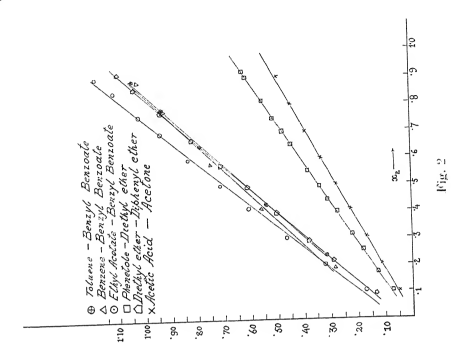
$$\log \eta_{sol} - \frac{1}{3} \log \sigma_{sol} = C_1 + C_2 x_2$$
.

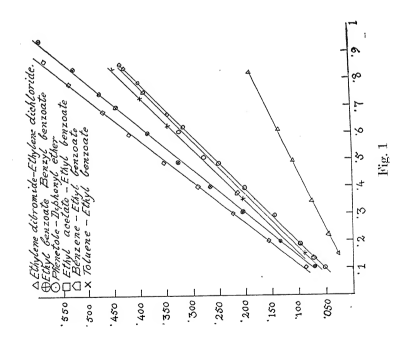
We have tried to apply the above equation to a number of binary systems and we have found the plot of  $\log \eta_{sol} - \frac{1}{2} \log \rho_{sol}$  against  $x_2$  to be generally a straight line. For convenience of plotting the values for the relative viscosities and relative densities have been used. The following eighteen systems have been analysed:—

(1) Phenetole-diphenyl ether, (2) ethyl benzoate-benzyl benzoate, (3) ethyl acetate-ethyl benzoate, (4) phenetole-diethyl ether, (5) diethyl ether-diphenyl ether, (6) ethyl acetate-benzyl benzoate, (7) benzene-ethyl benzoate, (8) toluene-ethyl benzoate, (9) toluene-benzyl benzoate, (10) benzene-benzyl benzoate, (11) ethylene dichloride-ethylene dibromide, (12) trichloracetic acid-acetone, (13) acetic acid-acetone, (14) phenol-benzene, (15) phenol-acetone, (16) trichloracetic acid-acetic acid-acetic acid-acetic acid-acetic acid-ethyl benzoate, (18) acetic acid-ethyl acetate.

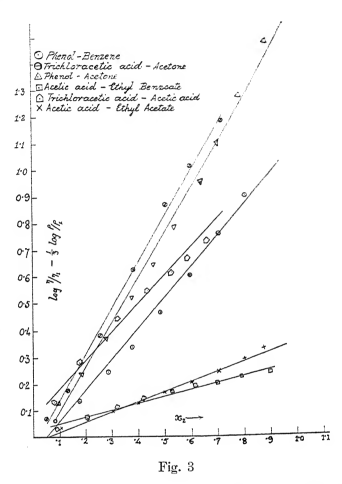
The first ten of the above-named systems have been analysed by Kendall and co-workers, whilst the system ethylene dichloride-ethylene dibromide has been investigated by MacFarlane and Wright. In every case these authors have found all previously proposed equations to be inadequate. On plotting  $\log \eta_{\rm sol} - \frac{1}{2} \log \rho_{\rm sol}$  against  $x_2$  we have found the graphs in the majority of cases to be very nearly straight lines. Thus the logarithmic equation represents the experimental values fairly satisfactorily. The graphs are given in figures 1 to 3 and the viscosity values are given in tables 1 to 18.

An expression for the viscosity of binary systems has been deduced by  $\operatorname{MacLeod^{10}}$  on the basis of the change of free space on mixing. He used the equation  $\eta_{\text{sol}} = \eta_1 \frac{m_1}{x} + \eta_2 \frac{m_2}{x} \frac{x_2}{x}$  where  $m_1$  and  $m_2$  are the molar fraction of the components and  $x_1, x_2, x_3$  are the amounts of free space of the two components and of the mixture respectively. Some of the systems analysed in the present paper have also been studied by MacLeod. It was, therefore, considered interesting to compare MacLeod's values with those obtained with the logarithmic equation. In the last three columns of tables 1 to 6 are given the percentage deviations obtained on the basis of MacLeod's equation and those calculated from the present equation. As will be seen from these tables, except in the case of the system ethyl acetate-benzyl benzoate, the logarithmic equation gives somewhat better values.





From an examination of the graphs (fig. 5) it will be seen that in the case of the systems phenol-benzene, trichloracetic acid-acetone, phenol-acetene, trichlor-



acetic acid-acetic acid, acetic acid-ethyl benzoate, and acetic acid-ethyl acetate the agreement is not so satisfactory as in the case of the remaining systems. The order of the deviations is too large to be due to any experimental error. In the case of these systems, however, one or both of the components have Polar properties and the nature of the components are dissimilar. There is thus a pronounced possibility of the formation of loose compounds. It will be reasonable to assume that in these cases the divergences are due to a form of loose chemical combination. The system phenol-benzene has also been studied by MacLeod. As will be seen from table 14, the divergences from both MacLeod's equation and the present equation are of the same order.

Thus the logarithmic equation is applicable to a large number of the systems named above. It would appear, therefore, that Andrade's viscosity equation applies fairly well to the case of binary solutions also. In applying the equation  $\log \eta_{\rm sol} - \frac{1}{3}\log \rho_{\rm sol} = C_1 + C_2$   $x_2$  it has been assumed that the constant A of Andrade's equation does not vary much with changes of concentration. That the above assumption is at least approximately true is borne out by the fact that in the case of a large number of the systems viscosity values are in accordance with the above equation.

In conclusion it might be stated that an expression similar to Andrade's equation has been deduced by MacLeod from his own equation.

One of us is indebted to the Patna University for a research scholarship which has enabled him to carry on the work.

 $\begin{tabular}{ll} Table & I \\ \hline \end{tabular}$  Phenetole-Diphenyl Ether

Mol fraction ether	$ ho_{ m obs}$	nobs	$\log \eta/\eta_1 - \frac{1}{3} \log \rho/\rho_1$	η <sub>cal</sub>	% age deviation	η from MacLeod equation	% age deviation
.0000	0.9618	.01158				•••	٧.,
<b>.</b> 099 <b>4</b>	0.9755	.01309	.05120	.01309	0.0	'01328	+1.5
·1873	0.9870	.01451	09425	.01451	0.0	01479	+1.5
<b>·</b> 2963	0.9993	.01632	<b>.</b> 14356	01645	+0.8	'01672	+2.4
.3998	1 0111	.01862	19907	01874	+0.6	.01883	+21
4895	1.0206	<b>.</b> 02096	24921	'02088	-0.4	.02092	-0.5
·6269	1.0346	02455	·31583	'02458	+0.1	02450	-0.5
6747	1.0396	02630	<b>3</b> 4503	02612	-0.7	.02610	-0.7
7928	1.0510	.03003	'40106	.03003	0.0	.02960	+0.3
<b>.</b> 8633	1.0573	03264	43641	.03264	0.0	.03230	+0.4
	:	į					

Table II

Ethyl Benzoate — Benzyl Benzoate

Mol fraction ethylbenzoate	$ ho_{ m obs}$	$\eta_{ m obs}$	$\log \eta_1 - \frac{1}{3} \log \rho_{\beta_1}$	$\eta_{cal}$	% age deviation	η from MacLeod equation	% age deviation
1037 1 1977 1 3094 1 4059 1 4946 1 6055 1 7027 7 7525 8451	10431 10524 10603 10683 10683 10760 10825 10910 10967 11005 11053 11104	02014 02371 02749 03249 03740 04309 04948 05695 06108 06898 08039		02995 02749 08282 08740 04286 04948 05695 06108 06995	+10 00 -05 00 -18 00 00 00 +14 +11	02381 02755 03245 03739 04265 05110 05930 06340 07100 08280	+04 +02 -01 00 -10 +31 +33 +36 +29 +30

 $\begin{tabular}{ll} Table & III \\ \hline {\bf Ethyl Acetate-Ethyl Benzoate} \end{tabular}$ 

Mol fraction nethyl-benzoate	$ ho_{ m obs}$	ηοbs	$\log^{\eta/\eta_1} - \frac{1}{3} \log \rho/\rho_1$	Heal	% age deviation	η from MacLeod equation	% nge deviation
10000 1008 2023 3011 4002 4956 6027 6869 7912 8784	0°8948 0°9204 0°9440 0°9598 0°9740 0°9866 0°9992 1°0101 1°0223 1°0320	004239 005188 006178 007247 008478 009851 01157 01292 01514 01704	108366 15585 12275 12875 135208 142005 146647 153358 158355	005263 006188 006188 007235 008406 009748 01149 01292 01529 01739	$\begin{array}{c} \dots \\ +1.4 \\ +0.2 \\ -0.1 \\ -0.85 \\ -1.20 \\ -0.7 \\ 0.0 \\ +1.0 \\ +1.5 \end{array}$	 '005240 '006455 '007470 '008690 '009760 '01140 '01280 '01492 '01708	+10 +42 +30 +11 -10 -14 -07 -16 -02

 $\begin{tabular}{ll} \it Table \ \it IV \end{tabular}$  Phenetole-Diethyl Ether

Mol fraction phenetole	$ ho_{ m obs}$	η̄obs	$\log \eta/\eta_1 - \frac{1}{3}\log \rho/\rho_1$	η <sub>eal</sub>	% age deviation	η from MacLeod equation	% age deviation
0000 0996 1697 2515 3118 3952 4421 4894 5499 6484 6974 7477 8145 9025 9255	7139 7458 7656 7891 8059 8282 8387 8488 8628 8853 8970 9083 9230 9427 9471	002233 002674 003057 003494 003866 004406 005154 005713 006667 007152 007856 008671 009993 01027	 '07196 '12630 '17995 '22084 '27366 '30985 '33821 '38057 '44390 '47250 '51146 '55200 '61058 '62177	002692 003050 003478 003841 004406 004749 005154 005674 006660 007162 007812 008665 009993 01039	$\begin{array}{c} \\ +0.7 \\ -0.2 \\ -0.5 \\ -0.6 \\ 0.0 \\ -1.2 \\ 0.0 \\ -0.7 \\ -0.1 \\ +0.1 \\ -0.6 \\ -0.1 \\ 0.0 \\ +1.1 \\ \end{array}$		+1.9 +1.2 +2.1 +2.5 +2.1 +0.7 -0.1 -2.2 -2.0 -1.4 -0.9 -1.9 -0.8 0.0

 $\label{eq:Table-V} Table\ V$  Diethyl Ether-Diphenyl Ether

Mol fraction phenyl ether	$ ho_{ m obs}$	nobs	$\log \eta/\eta_1 - \frac{1}{3} \log \ell/\rho_1$	neal	% age devia- tion	η from MacLeod equation	% age devia- tion
'0000 '0908 '2174 '2912 '3924 '4898 '5755 '6703 '7807 '8682 '9296	0.7139 0.7601 0.8213 0.8554 0.8974 0.9311 0.9583 0.9862 1.0181 1.0387 1.0542	002233 003106 004552 005733 007614 009926 01258 01631 02153 02737 03158	13353 28901 38032 49959 60945 70813 81682 93283 103411 109406	003183 004552 005639 007603 009827 01246 01614 02153 02741 03251	-1.0 -0.0 -1.0 -0.9 -1.0 0.0 +1.5 +2.9	'003061 '004523 '005738 '007735 '009935 '01227 '01570 '02110 '02655 '03153	-1.5 -0.5 +0.1 +0.7 +0.1 -2.5 -3.8 -2.0 -3.0 -0.2

Table VI Ethyl Acetate-Benzyl Benzoate

Mol fraction Benzylbenzoate	Ŋobs	$\log \eta/\eta_1 - \frac{1}{3} \log \rho/\rho_1$	મેહા]	age devia- tion	η from MacLeod equation	% age devia- tion
'0000       0'8948         '1020       0'9394         '2010       0'9720         '3027       0'9988         '4137       1'0247         '5000       1'0454         '5999       1'0625         '6998       1'0778         '7653       1'0867         '8574       1'0975         '9124       1'1048	004239 006406 009116 01275 01789 02299 03047 03881 04711 05938 07003	17228 32056 46232 60574 71179 88175 98475 101772 111679 118758	'006672 '009000 '01211 '01706 '02197 '02966 '03927 '04711 '06140 '07413	$\begin{array}{c} +41\\ -19\\ -50\\ -47\\ -41\\ -26\\ +12\\ +00\\ +34\\ \div55 \end{array}$	006458 006705 01240 01740 02298 03021 04010 04750 05984 07180	+0.9 -0.5 -2.5 -3.0 -0.1 -0.9 +0.9 -0.1 +2.5

Table VII
Benzene-Ethyl Benzoate

Mol fraction ester	$ ho_{ m obs}$	ηobs	$\frac{\log \eta_{\eta_1} - \frac{1}{3} \log \rho_{i \rho_1}}{-\frac{1}{3} \log \rho_{i \rho_1}}$
'0000 '1302 '3808 '5128 '6111 '7598 '8479	0'87809 0'90592 0'95700 0'98109 0'99387 1'01459 1'02350	**006051 **007244 **01018 **01200 **01327 **01557 **01709	07280 21258 27662 32227 38870 42787

Table VIII
Toluene-Ethyl Benzoate

Mol fraction ester	$ ho_{ m obs}$	$\eta_{ m obs}$	$\log \eta_{\eta_1}^{-\frac{1}{3}} \log \rho/\rho_1$
70000 71524 73546 76299 7349 78442	0°86244 0°89770 0°93921 0°98844 1°00530 1°03160	**005520 **006845 **009076 **01279 **01452 **01655	'08764 '20360 '34519 '39782 '45092

 $\begin{tabular}{ll} Table & IX \\ Toluene-Benzyl & Benzoate \\ \end{tabular}$ 

Mol fraction ester	$ ho_{ m obs}$	Nobs	$\log \eta/_{\eta_1} - \frac{1}{3} \log \rho/_{\rho_1}$
.0000	0°86244	`005520	0.31665
.2367	0°95282	`01183	
4261	1.0069 $1.0584$	.02015	0.53990
6502		.03614	0.78658
7890 9002	1.0829 $1.0966$	.05080 .06660	$0.93096 \\ 1.04664$

 $\label{eq:Table} Table \ X$  Benzene-Benzyl Benzoate

Mol fraction ester	$ ho_{ m obs}$	ηοbs	$\int \log \eta/_{\eta_1} - \frac{1}{3} \log \rho _{\rho_1}$
0000 1886 4124 5832 7827 8952	0°87309 0°95327 1°0175 1°0544 1°0831 1°1061	*006044 *01196 *02301 *03584 *05478 *06883	0.28372 0.55843 0.74471 0.92609 1.02220

 ${\it Table~XI}$  Ethylene Dichloride-Ethylene Dibromide

Mol fraction dibromide	$ ho_{ m obs}$	$\eta_{ m obs}$	$\log \eta/_{\eta_1} - \frac{1}{3} \log \rho/_{\rho_1}$
*000() *1499	1.238 1.389	.007812 .008648	···· •02755
·2209 ·3450	1.456 1.577	*009179 *01006	02755 04657 07475
·4941 ·6125	1.715 1.826	01117 01216	10809
*8250	2.016	01421	19010

Table XII
Trichloracetic acid-Acetone

Mol fraction acid	$ ho_{ m obs}$	$\eta_{\mathrm{obs}}$	$\log \eta_{1/\eta_1} - \frac{1}{3} \log \rho/\rho_1$
.0000	0.1875	003065	
.0484	0.8541	1003680	0.00223
.1316	0.0345	*004855	0.17169
2543	1.0230	.008156	0.38050
*3826	1.2090	01433	0.62319
5048	1.3190	02571	0.86764
5971	1.4000	03829	1.01363
7175	1.4830	05808	1.18640

Table XIII
Acetic acid-Acetone

Mol fraction acid	$ ho_{ m obs}$	$\eta_{ m obs}$	$\log\eta/_{\eta_1} - \tfrac13\log\rho/_{\rho_1}$		
.0000	0.7872	.003062	* * *		
.0996	0.8088	.003496	04731		
2035	0.8351	004046	11219		
3025	0.8568	004636	16751		
4049	0.8847	005350	22513		
<b>:49</b> 86	0.9064	.006098	27837		
5978	0.3333	.006994	33363		
·6968	0.9609	'008026	'38920		
·8015	0.9907	·009213	44489		
<b>.</b> 9037	1.0255	.01036	49081		

Table XIV
Phenol-Benzene

Mol frac- tion p henol	$ ho_{ m obs}$	$\eta_{ m obs}$	$\log \eta/\eta_1 - \frac{1}{3} \log \rho/\rho_1$	$\eta_{\mathrm{cal}}$	% age deviation	η <sub>cal</sub> from MacLeod Equation	% age devia- tion
.0000	0.8772	100629	•••				
.0506	0.888.0	.00683	.03400				
.0803	0.8949	.00724	05820				
1720	0.9133	.00862	13253				
2845	0.9370	'01126	24334	'01211	+7.5	'0123	+8.6
37575	0 9549	.01401	*33550				
·4836	0.9766	.01911	46492	.02108	+10.3	.0196	+2.0
:5923	0.9976	.02645	.60466				
7037	1.0194	03811	76061	03771	-0.3	.0349	+S.4
.8043	1.0383	.02320	90528			A service	

 $\begin{array}{c} \textit{Table XV} \\ \textit{Phenol-Acetone} \end{array}$ 

Mol fraction phenol	$ ho_{ m obs}$	$\eta_{ m obs}$	$\log \eta/_{\eta_1} - \frac{1}{3} \log \rho/_{\varrho_1}$
**************************************	0.8031	00360	0.12836
	0.8425	00486	0.28368
	0.8768	00635	0.28368
	0.9085	00868	0.36445
	0.9406	01256	0.51995
	0.9642	01688	0.64495
	0.9851	02358	0.78679
	1.0090	03670	0.95585
	1.0237	04950	1.10824
	1.0420	07480	1.27988
	1.0628	11930	1.47998

 $Table\ X\,VI$  Trichloracetic acid-Acetic acid

Mol fraction trichloracetic acid	ρ <sub>obs</sub>	ηοbs	$\log \eta/\eta_1 - \frac{1}{3} \log \rho/\rho_1$
0000 0737 1777 3209 4348 5262 5853 6581	1'049 1'129 1'223 1'337 1'409 1'457 1'491 1'508	01121 01532 02228 03362 04346 05176 05859 06854	12483 127592 144191 154595 161682 166752 173383

 $\begin{tabular}{ll} Table & XVII \\ Acetic acid-Ethyl & Benzoate \\ \end{tabular}$ 

Mol fraction ester	$ ho_{ m obs}$	$\eta_{ m obs}$	$\log \eta/_{\eta_1} - \frac{1}{3} \log \rho/_{\rho_1}$
'0000	1.050	01121	
'0868	1.049	01202	'03036
'2044	1.049	01322	'07196
'3174	1.048	01446	'11051
'4178	1.048	01538	'13771
'5250	1.047	01651	'16829
'6118	1.047	01727	'18808
'6955	1.047	01797	'19871
'7871	1.046	01874	'22336
'8959	1.046	01948	'24078

Table XVIII

Acetic acid-Ethyl Acetate

Mol fraction acid	$ ho_{ m obs}$	$\eta_{ m obs}$	$\log \eta_{\eta_1} - \frac{1}{2} \log \rho_{\eta_1}$
.0000	0.8948	:	
1049	0.9092	004590	03234
2070	0.9211	004949	706337
.3037	0.9308	005331	09396
18990	0.9417	005762	12621
4985	0.9557	006289	16224
5996	0.9697	006890	19962
·6988	0.9850	007668	24376
8011	1.0015	008590	29067
8742	1.0165	009430	32915

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## ACTION OF PARA-TOLUENE-SULPHONYL CHLORIDE ON PHENOLS CONTAINING AZO GROUPS

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#### SUMMARY

- 1. The action of para-toluene-sulphonyl chloride on some phenols containing azo groups has been studied. In each case an ester has been obtained in the presence of diethylaniline as the condensing reagent.
- 2. Monoazo and disazo phenols yield esters easily. Trisazophenol does not form an ester. In no case is the OH group replaced by Cl atom.
- 3. The reactivity of these esters with aniline has also been studied and it has been found that the esters are reactive only when at least one  $(NO_2)$  group is present in the ortho position nearest to the OH group, and another  $NO_2$  or CH<sub>3</sub> group in the second benzene ring.

In 1908 Ullmann and Nadai<sup>c-8</sup> observed that para-toluene-sulphonyl chloride reacts with 2:4-dinitro-phenol in two ways. In presence of sodium carbonate as condensing reagent, it forms a para-toluene-sulphonyl ester, but if diethylaniline is used as a condensing reagent, the -OH group is replaced by chlorine atom and 1:2:4-chloro-dinitro-benzene is obtained. Mono-nitro-phenols, however, yield esters in presence of sodium carbonate as well as diethylaniline. This reaction was studied by Ullmann and his collaborators<sup>9</sup> and later more extensively by Sane and Joshi, <sup>1-3</sup> who investigated the influence of the various groups vix., -CH<sub>3</sub>, C1, Br, I, -NO<sub>2</sub> etc., on the replaceability of the -OH group in this reaction.

The chloro derivatives which are formed by this reaction contain an active chlorine atom.<sup>4, 5</sup> The para-toluene-sulphonyl esters of dinitrophenols which yield chloro-dinitro-benzenes in presence of diethyl aniline are also reactive and when heated with aniline react as follows:—

All para-toluene-sulphonyl esters of phenols do not react in the above manner, and this reactivity is found to depend upon the groups which are present in the phenols.

In the following paper the influence of diazo groups alone or along with NO<sub>2</sub> and Br groups has been examined. Mono and disazo-phenols form ester easily but the -OH group is not replaced by chlorine atom, when diethylaniline is used as a condensing reagent. Symmetrical trisazophenol does not form an ester. The esters formed are, however, reactive, *i.e.*, react with aniline in the manner shown above, only when at least one NO<sub>2</sub> (acid) group is present, in the ortho position nearest to the -OH group; but one NO<sub>2</sub> is not sufficient by itself thus:—

does not react with aniline but when NO<sub>2</sub> or CH<sub>3</sub> group is introduced into the second benzene ring the ester becomes reactive, thus:—

1. 
$$NO_2$$
,  $C_6H_6$ ,  $N; N, C_6H_5$ ,  $NO_2$ ,  $O.SO_2$ ,  $C_6H_4$ ,  $CH_3 + NH_2$ ,  $C_6H_5 = 4$  11' 3' 4' 4'  $CH_3 + NO_2$ ,  $C_6H_4$ ,  $CH_5 + NO_2$ ,  $C_6H_4$ ,  $N; N, C_6H_5$ ,  $NO_2$ ,  $NH, C_6H_5$  2.  $CH_3$ ,  $C_6H_4$ ,  $N; N, C_6H_5$ ,  $NO_2$ ,  $C_6H_4$ ,  $N; N, C_6H_5$ ,  $NO_2$ ,  $C_6H_4$ ,  $CH_3 + NN_2$ ,  $C_6H_5 = 4$   $CH_3$ ,  $C_6H_4$ ,  $CG_4H_5$ ,  $CG_5H_5$ ,  $CG_6H_6$ ,

#### EXPERIMENTAL

The toluene-sulphonyl esters of the azo-nitrophenols were obtained by heating calculated quantities of the respective azo-nitrophenols, paratoluene-sulphonyl-chloride and diethylaniline on a water-bath for 4 hours. The mixture was cooled and then treated with dilute HC1, in order to decompose the unchanged diethylaniline. The aqueous liquid was then decanted off and the residue shaken with a little alcohol when in most cases the solid ester separated out. This was filtered and recrystallised from a suitable solvent. The solvent commonly used was glacial acetic acid and in some cases toluene.

The reactivity of the above esters with aniline was also studied separately in each case, and it was observed that esters were reactive, only when at least one NO<sub>2</sub> group is present in the ortho position nearest to the OH group, and another NO<sub>2</sub> or CH<sub>3</sub> group in the second benzene ring—yielding substituted diphenylamine compounds.

For the preparation of the diphenylamine compounds, calculated quantities of the toluene-sulphonyl-esters of the azo-nitrophenol, freshly distilled aniline, ethyl alcohol and fused anhydrous sodium acetate were refluxed for an hour over a water-bath. The alcohol was then distilled off and the excess of aniline removed by HCl. The solid mass, which separated out was thoroughly washed with water and finally recrystallised from acetic acid or toluene. The compounds obtained are generally deeply coloured, either deep red, orange or yellow.

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Melting
1'  1  3:5  4  4. N: N. C <sub>6</sub> H <sub>3</sub> . NO <sub>2</sub> . X
$132^{\circ}$ C
154°C
125°C
$135^{\circ}$ C
124°C
1'  1  3  4  1
$180^{\circ}$ C
$163^{\circ}\mathrm{C}$
178°C
178°C
- 1

The action of  $\alpha$ -naphthylamine was also studied on one ester. In this case 2 gms, of the ester (NO<sub>2</sub> C<sub>6</sub>H<sub>4</sub>, N: N. C<sub>6</sub>H<sub>5</sub>, NO<sub>2</sub>, O.SO<sub>2</sub>, C<sub>6</sub>H<sub>4</sub>, CH<sub>5</sub>) was dissolved in about 40 c.e. boiling amyl alcohol, and then about 2 gms, of  $\alpha$ -naphthylamine and 2 gms, of fused sodium acetate added. This was then heated for an hour under a reflux condenser. On cooling a solid mass separated out which was filtered and then treated with HCl and hot water in order to remove the unchanged  $\alpha$ -naphthylamine, the residue was then recrystallised from toluene.

The table on page 220 summarises the experimental results of the present investigation.

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### IONISATION OF F-REGION BEFORE SUNRISE

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#### SUMMARY

Curves have been drawn for the time of sunrise and sunset at different heights for Allahabad for the whole year and with the help of these curves, the height where the sun's rays are falling at the time of minimum ionisation of the F-region just before sunrise is found. Similar calculations have been made for Washington and Slough and from these it has been shown that this height is greater in winter and smaller in summer for all the latitudes. For higher latitudes (e.g., Slough) the ionization always begins to rise after sunrise in the F-region while for lower latitudes (e.g., Washington and Allahabad) we can roughly say that the ionization begins to rise before sunrise in winter and after sunrise in summer.

It is now well recognised that the ionisation of the upper atmosphere is due to the action of the ultra-violet rays of the sun on the molecules of N<sub>2</sub> and O<sub>2</sub>. The correctness of the solar origin of ionisation has been established in the eclipse expeditions of 1932 in the case of the E and F1-layers, but in the case of F2-layer, the observations have not yet yielded any decisive result. The F2 layer has been found to behave in an anomalous way in other respects as well. Pederson showed on the basis of the solar origin of ionisation that in the daytime the maximum concentration of electrons should vary as  $\cos^{\frac{1}{4}}\psi$  (where  $\psi$ =zenith distance of the sun), but this law is apparently verified only in the case of the E and F1-layers and not in F2-layer. The F-layer2 in the Ionisphere breaks up into two regions F1 and  $\mathrm{F}_2$  during the day when the sun is fairly high and its angular zenith distance is less than about 40°. These two layers coalesce later, toward night fall and we have only one region which is usually supposed to be a continuation of the F<sub>2</sub>-region. The  $F_2$ -region is thus only a daylight phenomenon. The ionization in lower  $F_1$ layer is always found to have a maximum value near about noon as we should expect if the ionization be due to solar radiation but it is not so with the F2-region. The daytime ionization in the F2-region is found to have two maxima in the summer, one a few hours before noon and the other late in the afternoon.<sup>5</sup> The ionisation in the F2-layer during the night is found to have one minimum during the summer and two minima in the winter, and there occurs a maximum at about 1 A.M. To explain these anomalous behaviours of the F2-region, Appleton thinks that during daytime in summer the F2-layer heights are heated to about 1200° K by sunlight and then the ionised layers expand and concentration falls. So there is a minimum at noon. The very hypothesis of a higher temperature on a summer morning than on a winter morning also explains the anomaly of two minima in the winter and one minimum in summer at night as a reduced ion production due to the reduced air density at the level of maximum ion production. But Berkner<sup>3</sup> contended that if this explanation were correct then in the southern hemisphere, as the seasons are reversed the conditions in the layer ought to be reversed, but his observations showed that this was not the case. The variation of the maximum density curve was found to be the same in both the hemispheres at the same instant throughout the year. It appears, therefore, that the  $F_2$ -region ionisation is a complex phenomenon. A minimum of ionisation in the region has however been always observed sometime before sunrise. The present investigation was undertaken to see whether the increase in ionisation begins to take place when the sun rises in the  $F_2$ -region.

#### SUNRISE AND SUNSET IN THE UPPER REGIONS

For this we require a formula to calculate the time of sunrise and sunset at different heights for different seasons. The height 'h' at which the sun rises when the time is 'l' is given by

$$h = 6480 \text{ (sec}\eta - 1)$$
 ... (1)

where  $\sin \varphi = -(\cos t' + \tan \delta \tan \varphi) \cos \delta \cos \varphi$  ... (2)

t'=time between the time of sunrise at height 'h' and midday. (This will give us apparent time. To convert it into mean solar time, the equation of time is to be added.)

 $\eta$ =The angle by which the sun is below the horizon at the time t'

 $\varphi$ =Latitude of the place

 $\delta$  = Declination of the sun.

The calculations for the time of sunrise and sunset at different heights over Allahabad were made with the aid of formulæ (1) and (2) for the whole year at intervals of fifteen days and these are represented in figs. 1, 2, 3 and 4. Figs. 1 and 2 are for sunrise, while figs. 3 and 4 are for sunset. From these curves we can at once know the time of sunrise and sunset at any height on any day of the year.

At Allahabad [Lat. 25° 25′ 25″] observations of penetration frequency for the F-layers were taken continuously for twentyfour hours in the months of October, November and December 1936, once a week and curves were drawn between the penetration frequency and the Indian standard time. These observations have been reported elsewhere by Pant and Bajpai. From the curves given in their paper the time of minimum ionisation can be easily found out. Corresponding to this time of minimum ionisation the height at which sun's rays were falling can be read off from the curves in figs. 1 and 3. These are shown in the first part of table I. Similar tables have been made from data taken by Gilliland<sup>4</sup> for Washington [Lat. 39° 2′ 0″ N] and by Appleton<sup>4</sup> and Naismith for Slough [Lat. 51° 29′ 30″ N]. These tables form part 2 and part 3 of table I.

 $Table\ I$ 

Part	Place	Date	Time of ground sunrise	T <sub>1</sub> Time of first mini-ionisation	Sunrise at height corresponding to T <sub>1</sub> in kms.	T <sub>2</sub> Time of second mini-ionisation	Height of sunrise corresponding to T <sub>2</sub> in kms.	Whether minimum before or after sunriseat I'-layer
1	Allahabad [25° 25′ 25′]	3-10-36 18-10-36 15-11-36 21-11-36	0553 0559 0617 0621	0400 0400 0430 0530	690 735 580 135	0030	 56560	Before ,, After
•)	Washington [39° 2′ 0″]	May, 1933 June, 1933 July, 1933 August, 1933 September, 1933 October, 1933 November, 1933 December, 1934 January, 1934 February, 1934 March, 1934 April, 1934	0448 0433 0440 0511 0540 0609 0642 0711 0719 0653 0613 0525	0400 0400 0400 0200 0430 0430 0600 0600 0600 0510 0410 0400	76 35 45 980 175 358 58 159 208 403 603 246	2300 2300 2300 2300 2320 2400 	6030 10500 7193 12310 8740	After "Before After Before After "Before "Before" "Before"
3	Slough [51° 29′ 30°]	15-1-34 15-2-34 15-3-34 15-4-34 15-6-34 15-7-34 15-8-34 15-9-34 15-10-34	0800 0717 0617 0507 0410 0342 0405 0445 0532 0630 0717	0726 0615 0526 0430 0335 0316 0340 0400 0455 0535 0615	24 98 69 9 23 11 7 38 42 97 88			After "" "" "" "" "" "" "" "" "" "" ""

In table I the places for which the calculations have been made are shown in the second column. The dates for which calculations have been made are shown in column third. The time of ground sunrise is shown in column fourth. The time of first minimum ionisation just before sunrise is shown in column fifth. The height at which the sun rises at the time shown in column fifth is shown in column sixth. The time of second minimum ionisation before sunrise is shown in column seventh and the height at which the sun rises at the time given in column seventh, is shown in column eighth, while the ninth column shows whether the ionisation begins to rise before or after the sun rise in the  $F_2$ -region.

In the case of Allahabad and Slough the data are for particular day while for Washington they are the average of the whole month.

The data for Allahabad show that on 3rd October, 18th October, and 15th November, 1936, the ionisation begins to rise before sunrise at the F-region, while on the 21st November, 1936 it begins after the sunrise at the F-region. There is also a second minimum at 0030 on the same day. For Washington we see that in May, June, July, September, November and December, 1933 as well as in January 1934 the ionisation begins to rise after sunrise in the F-region, while in August and October, 1933 and February, March and April, 1934 it begins to rise before sunrise in the F-region. There is also a second minimum in October, November, December, January and February, always lying between 2300 and 2400. For Slough the ionisation begins to rise always after sunrise. However it is apparent from the data that in winter the height at which the sun is rising at the time of minimum ionisation is greater than in summer.

#### Conclusion

The data on the whole show that the height corresponding to the time of minimum ionisation is greater for winter than for summer for all latitudes. It also indicates that for higher latitudes, the ionisation in the  $F_2$ -region always begins to increase after the sun has risen in the regions from which the  $F_2$ -echoes are coming. This is evident from the data for Slough. For lower latitudes this is not so. For both Washington and Allahabad we find that at times the ionisation begins to increase before sunrise in the  $F_2$ -region, while at times the ionisation begins to increase after sunrise.

For lower latitudes we can roughly say that the ionisation begins to increase after sunrise in the  $F_2$ -region in summer and before sunrise in the  $F_2$ -region in winter.

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